

*Renumbering  
of claims  
Rule 1.126.*

5. The composition according to claim 1 wherein the amount of S-tofisopam or a prodrug, or pharmaceutically acceptable salt thereof is 99% or more by weight of the total weight of tofisopam.

*Renumbered  
under Rule  
1.126*  
Claim 6-<sup>27</sup>~~28~~ (cancelled)

Claim <sup>28</sup>~~29~~ (added)

<sup>28</sup>~~29~~. (Added) A composition according to claim 1, wherein the composition is for intraperitoneal, subcutaneous, intranasal, intramuscular, intrathecal, sublingual, rectal, intravenous infusion, transdermal delivery or oral administration.

<sup>29</sup>  
Claim <sup>30</sup>~~31~~ (added)

<sup>29</sup>  
<sup>30</sup>. (Added). A composition according to claim 1, wherein the amount of S-tofisopam, prodrug, or a pharmaceutically acceptable salt thereof is from approximately 10 mg to 1200 mg.

<sup>30</sup>  
Claim <sup>31</sup>~~32~~ (added)

<sup>30</sup>  
<sup>31</sup>. (Added) A composition according to claim 1, wherein the amount of S-tofisopam, prodrug or pharmaceutically acceptable salt thereof is from approximately 50 mg to 600 mg.

<sup>31</sup>  
Claim <sup>32</sup>~~33~~ (added)

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~~32~~. (Added) A composition according to claim 1, wherein the amount of S-tofisopam, prodrug or pharmaceutically acceptable salt thereof is from approximately 100 mg to 400 mg.

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Claim ~~33~~ (added)

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~~33~~. (Added) A composition according to claim 1, wherein the amount of S-tofisopam, pro-drug or pharmaceutically acceptable salt administered is less than 30 mg/kg.